518 Recapcipio o 2 Aug 2001

FORM PT (REV. 11-	D-1390 U.S. DEPARTMENT OF COM 2000)	MMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY'S DOCKET NUMBER					
*		TO THE UNITED STATES	Mo-6479/LeA 33.510_					
		ED OFFICE (DO/EO/US)	U.S. APPLICATION NO. de kowa see 37-GFR 1.5					
TA TOP DE		IG UNDER 35 U.S.C. 371	To be Assigned					
	RNATIONAL APPLICATION NO.	INTERNATIONAL FILING DATE 26 January 2000 (26.01.00)	PRIORITY DATE CLAIMED 05 February 1999 (5.02.99)					
	EP00/00576 OF INVENTION METHOD FOR	HYDROGENATING AROMATIC POL	<u> </u>					
SPECI	AL CATALYSTS							
APPLI	CANT(S) FOR DO/EO/US 1) VOIK	ker Wege; 2) Konstadinos Douzinas; 3) Jo	ohann Rechner					
Applic	ant herewith submits to the United St	ates Designated/Elected Office (DO/EO/US)	the following items and other information:					
1. X	This is a FIRST submission of items	s concerning a filing under 35 U.S.C. 371.						
2.	This is a SECOND or SUBSEQUE	NT submission of items concerning a filing un	nder 35 U.S.C. 371.					
3. X		national examination procedures (35 U.S.C. 3	371(f)). The submission must include					
4. X	items (5), (6), (9) and (21) indicated The US has been elected by the expi		article 31).					
5. X	The US has been elected by the expiration of 19 months from the priority date (Article 31). A copy of the International Application as filed (35 U.S.C. 371(c)(2))							
		d only if not communicated by the Internatio	onal Bureau).					
	b. has been communicated by	y the International Bureau. lication was filed in the United States Receivii	ng Office (RO/LIS)					
6.X		he International Application as filed (35 U.S.)						
0.12	a. X is attached hereto.	international reprientation as med (55 5.2.	C. 371(c)(2)).					
	b. has been previously submit	tted under 35 U.S.C. 154(d)(4).						
7.	Amendments to the claims of the International Aplication under PCT Article 19 (35 U.S.C. 371(c)(3))							
		ed only if not communicated by the Internation	onal Bureau).					
	<u> </u>	by the International Bureau.	the NOT ownered					
	=	ver, the time limit for making such amendmen	nts has NOT expired.					
۵ □	d. In have not been made and wi		-1- 10 (25 H C C 271 (a)(2))					
8. 🔲		the amendments to the claims under PCT Artic	:16 19 (33 U.S.C. 371 (c)(3)).					
9. □	An oath or declaration of the inventor							
10.	An English lanugage translation of the Article 36 (35 U.S.C. 371(c)(5)).	he annexes of the International Preliminary E	xamination Report under PCT					
	ns 11 to 20 below concern documen							
11.	An Information Disclosure Stateme	ent under 37 CFR 1.97 and 1.98.						
12.	An assignment document for record	ding. A separate cover sheet in compliance v	with 37 CFR 3.28 and 3.31 is included.					
13. X	A FIRST preliminary amendment.		{					
14.	A SECOND or SUBSEQUENT pr	reliminary amendment.						
15.	A substitute specification.		*					
16.	A change of power of attorney and	i/or address letter.						
17.	A computer-readable form of the se	equence listing in accordance with PCT Rule	13ter.2 and 35 U.S.C. 1.821 - 1.825.					
18. 🗓	A second copy of the published int	ternational application under 35 U.S.C. 154(d))(4).					
19.	A second copy of the English lang	guage translation of the international application	on under 35 U.S.C. 154(d)(4).					
20. X	Other items or information:		·					
PTO F	orm 1449 and references listed the	erein; Preliminary Amendment w/Abstrac	et					
			·					

1647 Rec's PUT/PTO 10 2 Aug 2001 INTERNATIONAL APPLICATION NO. ATTORNEY'S DOCKET NUMBER Mo-6479/LeA 33,510 PCT/EP00/00576 CALCULATIONS PTO USE ONLY 21. X The following fees are submitted: BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)): Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a (2)) paid to USPTO and International Search Report not prepared by the EPO or JPO International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO\$710.00 International preliminary examination fee (37 CFR 1.482) paid to USPTO but all claims did not satisfy provisions of PCT Article 33(1)-(4) \$690.00 International preliminary examination fee (37 CFR 1.482) paid to USPTO and all claims satisfied provisions of PCT Article 33(1)-(4) \$100.00 **ENTER APPROPRIATE BASIC FEE AMOUNT** = 860.00 Surcharge of \$130.00 for furnishing the oath or declaration later than \$ months from the earliest claimed priority date (37 CFR 1.492(e)). NUMBER FILED **CLAIMS** NUMBER EXTRA \$ RATE Total claims -20 =x \$18.00 \$ 0 0.00 Independent claims 2 -3 =\$ x \$80.00 0.00 MULTIPLE DEPENDENT CLAIM(S) (if applicable) \$ + \$270.00 0.00 TOTAL OF ABOVE CALCULATIONS \$ 860.00 Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above \$ are reduced by 1/2. 0.00 \$ 860.00 Processing fee of \$130.00 for furnishing the English translation later than \$ months from the earliest claimed priority date (37 CFR 1.492(f)). TOTAL NATIONAL FEE 860.00 Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be \$ accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property 0.00 TOTAL FEES ENCLOSED = \$ Amount to be refunded: \$ charged: A check in the amount of \$ ______ to cover the above fees is enclosed. Please charge my Deposit Account No. 13-3848 in the amount of \$ 860.00 to cover the above fees. A duplicate copy of this sheet is enclosed. c. | X The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 13-3848 ... A duplicate copy of this sheet is enclosed. Fees are to be charged to a credit card. WARNING: Information on this form may become public. Credit card information should not be included on this form. Provide credit card information and authorization on PTO-2038. NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137 (a) or (b)) must be filed and granted to restore the application to pending status. SEND ALL CORRESPONDENCE TO: SIGNATURE Aron Preis 00157 NAME PATENT TRADEMARK OFFICE 29,426 REGISTRATION NUMBER

TRANSMITTAL LETTER TO THE UNITED STATES RECEIVING OFFICE

Date 2 August 2001

International Application No. PCT/EP00/00576

Attorney Docket No. Mo-6479/LeA 33 510

	TIED STATES RECEIVING OFFIC		Attorne	y Docket No.	Mo-64	479/LeA 33,510	7
I.	Certification under 37 CFR 1.10 (if ap	plicable)	- <u> </u>	0	9/891	073Z	_
	ET146886068US				2 Augu	ıst 2001	
	Express Mail mailing number					Deposit	\Box
!	I hereby certify that the application/correspondence attached he Addressee" service under 37 CFR 1.10 on the date indicated a	ereto is being above and is a	deposited with ddressed to As	the United St sistant Commi	ates Postal Servi ssioner for Pater	ice "Express Mail Post Office to nts, Washington, D.C. 20231.	
	Dans Vlater.	7			Donna .	J. Veatch	\neg
	Signature of person mailing correspondence			Typed or pr	inted name of p	erson mailing correspondence	
II.	X New International Application					Earliest priority date	=
	TITLE METHOD FOR HYDROGENATIN PRESENCE OF SPECIAL CATAL	IG AROM YSTS	ATIC POL	YMERS I	N THE	(Day/Mon/Year) (05/02/1999)	
•	application for purposes of determining whether other purposes, the following information is sup A. The invention disclosed was not made in the B. There is no prior U.S. application relating to the following prior U.S. application (s) con attached international application. (NOTE PCT/RO/101 (Request) and this listing documents.)	a license for opplied. (Note the United State of this inventation subject of priority to	foreign trans c: check as ma ates. ion. matter which these applic	mittal should any boxes as a is related to ations may	d and could be apply): o the invention or may not be	granted and for	
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	application no.		file	d on] [
III.	in paragraph C. above. The additional subject and DOES NOT ALTER MIC manner which would require the U.S. applied defense agencies under 35 U.S.C. 181 and 3 A Response to an Invitation from the RO A. A Request for An Extension of Time to B. A Power of Attorney (General or Regular Control of Contro	GHT BE CO cation to have 37 CFR 5.1. /US. The formula of the property of the second s	ONSIDERED ye been made See 37 CFR	TO ALTER available for 5.15	r inspection b	y the appropriate	
	C. Replacement pages:						
	7.0	equest (PCT/F	(O/101)	pages		of the figures	
		escription		pages		of the abstract	
	D. Submission of Priority Documents	laims					
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	E. Fees as specified on attached Fee Calculate	tion sheet for		of annex			
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IV.	A Request for Rectification under PCT 9	1 /	A Petition	A	Sequence L	isting Diskette	
v	X Other (please specify): PTO Form 1449 a	nd referen	ices listed t	herein; Pro	eliminary A	mendment w/Abstract	
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PATENT APPLICATION Mo-6479 LeA 33,510

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

APPLICATION	ON OF)
VOLKER W	EGE ET AL) PCT/EP 99/00576)
SERIAL NUI	MBER: TO BE ASSIGNED)
FILED:	HEREWITH	
TITLE:	METHOD FOR HYDROGENATING AROMATIC POLYMERS IN THE PRESENCE OF SPECIAL CATALYSTS	

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents Washington, D.C. 20231 Sir:

Prior to its examination kindly amend the English language translation of the enclosed application as follows:

"Express Mail" mailing label number	ET146886068US August 2, 2001
Date of Deposit or fee is	s being deposited with the United States
Postal Service "Express Mall Post Office 1.10 on the date indicated above and is of Patents and Trademarks, Washington	addressed to the Assistant Commissioner , D.C. 20231
Donna	J. Veatch
(Name of person)	mailing paper or fee)
Signature of person	mailing paper or fee)

Delete the Title of the application - appearing in pages 1 and 13 - and insert therefor:

-- Method for Hydrogenating Aromatic Polymers in the Presence of Special Catalysts --.

IN THE ABSTRACT:

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Please revise the Abstract of the disclosure at page 13 to read:

"A process for hydrogenating aromatic polymers is disclosed. The process entails using a supported catalyst wherein support contains alumina and where the catalyst comprise least one metal of subgroup VIII of the Periodic Table. The catalyst is characterized in that the pore volume of the pores having diameters of 1000 to 10,000 Angstrom is 15 to 100% relative to the total pore volume, as determined by mercury porosimetry."

IN THE CLAIMS:

Please amend the claims as follows:

- 5. A process according to Claim 1 wherein the catalysts have a pore volume, as measured by nitrogen sorption, of less than 10% for pore diameters less than 600 A.
- 6. A process according to Claim 1 wherein the metals are selected from nickel, platinum, ruthenium, rhodium and palladium.
- 7. A process according to Claim 1 wherein the process is conducted in the presence of solvents which can be used for hydrogenation reactions.
- 8. A process according to Claim 1 wherein the catalyst has a specific surface greater than 5 m²/g.

Please add the following:

--9. A process for hydrogenating aromatic polymers comprising using a supported catalyst wherein support contains alumina and where the catalyst comprise at least one metal of subgroup VIII of the Periodic Table, said catalyst characterized in that the pore volume of the pores having diameters of 1000 to 10,000 Angstrom, is 15 to 100% relative to the total pore volume, said pore volume determined by mercury porosimetry. --

REMARKS

The present amendment seeks to place the application in better conformance with U.S. practice. A page containing an Abstract of the disclosure is enclosed.

Attached hereto is a marked-up version of the changes made to the specification and claims by the current amendment. The attached page is titled "VERSION WITH MARKINGS TO SHOW CHANGES MADE"

Entry of the amendment is requested.

Respectfully submitted,

VOLKER WEGE KONSTADINOS XOUZINAS JOHANN RECHNER

Ву__

Aron Preis

Attorney for Applicants Reg. No. 29,426

Bayer Corporation 100 Bayer Road Pittsburgh, Pennsylvania 15205-9741 (412) 777-8341 FACSIMILE PHONE NUMBER: (412) 777-8363 s:\kgb\ap2409pa Mo-6479

Version with markings to show changes made

Delete the Title of the application - appearing in pages 1 and 13 - and insert therefor:

-- Method for Hydrogenating Aromatic Polymers in the Presence of Special Catalysts --.

IN THE ABSTRACT:

Please revise the Abstract of the disclosure at page 13 to read:

"A process for hydrogenating aromatic polymers is disclosed. The process entails using a supported catalyst wherein support contains alumina and where the catalyst comprise least one metal of subgroup VIII of the Periodic Table. The catalyst is characterized in that the pore volume of the pores having diameters of 1000 to 10,000 Angstrom is 15 to 100% relative to the total pore volume, as determined by mercury porosimetry."

Please amend the claims as follows

- 5. A process according to [one or more of the preceding claims,] <u>Claim 1</u> wherein the catalysts have a pore volume, as measured by nitrogen sorption, of less than 10% for pore diameters less than 600 A.
- 6. A process according to [one or more of the preceding claims,] <u>Claim 1</u> wherein the metals are selected from nickel, platinum, ruthenium, rhodium and palladium.
- 7. A process according to [one or more of the preceding claims,] Claim 1 wherein the process is conducted in the presence of solvents which can be used for hydrogenation reactions.
- 8. A process according to [one or more of the preceding claims,] Claim 1 wherein the catalyst has a specific surface greater than 5 m²/g.

Mo-6479 -4-

Please add the following:

--9. A process for hydrogenating aromatic polymers comprising using a supported catalyst wherein support contains alumina and where the catalyst comprise at least one metal of subgroup VIII of the Periodic Table, said catalyst characterized in that the pore volume of the pores having diameters of 1000 to 10,000 Angstrom, is 15 to 100% relative to the total pore volume, said pore volume determined by mercury porosimetry. --

LeA 33,510 – foreign countries

-13 -

METHOD FOR HYDROGENATING AROMATIC POLYMERS IN THE PRESENCE OF SPECIAL CATALYSTS

ABSTRACT OF THE DISCLOSURE

A process for hydrogenating aromatic polymers is disclosed. The process entails using a supported catalyst wherein support contains alumina and where the catalyst comprise at least one metal of subgroup VIII of the Periodic Table. The catalyst is characterized in that the pore volume of the pores having diameters of 1000 to 10,000 Angstrom is 15 to 100% relative to the total pore volume, as determined by mercury porosimetry.

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A process for the hydrogenation of aromatic polymers in the presence of special catalysts

The present invention relates to a process for the hydrogenation of aromatic polymers, which is characterised in that metals of subgroup VIII of the periodic table are present, together with a support comprising alumina. The catalysts have a special pore volume distribution. By this means, the aromatic units of aromatic polymers can be hydrogenated completely and without significant reduction of their molecular weights.

The hydrogenation of aromatic polymers is already known. DE-AS 1 131 885 describes the hydrogenation of polystyrene in the presence of catalysts and solvents. Aliphatic and cycloaliphatic hydrocarbons, ethers, alcohols and aromatic hydrocarbons are mentioned as solvents. A mixture of cyclohexane and tetrahydrofuran is stated to be preferred. Silica and alumina supports for the catalysts are mentioned in general, but the physicochemical structure thereof is not described.

EP-A-322 731 describes the production of what are predominantly syndiotactic polymers based on vinylcyclohexane, wherein a styrene-based polymer is hydrogenated in the presence of hydrogenation catalysts and solvents. Cycloaliphatic and aromatic hydrocarbons are mentioned as solvents, but ethers are not mentioned.

DE 196 24835 (= EP-A 814 098), which relates to the hydrogenation of polymers using ruthenium or palladium catalysts in which the active metal is deposited on a porous support, describes the hydrogenation of olefinic double bonds of polymers.

Aromatic regions are hydrogenated by less than 25 %, and in general are hydrogenated within a range from 0 to about 7 %. The choice of solvent is not critical.

It is also known (WO 96/34896 = US-A-5,612,422) that silica-supported catalysts for the hydrogenation of aromatic polymers, which catalysts have small pore diameters (200-500 Å) and large specific surfaces (100-500 m²/g), can result in incomplete hydrogenation and to a breakdown of the polymer chain. The use of special silica"Express Mail" mailing label number ET146886068US

Date of Deposit ______ August 2, 2001

I hereby certify that this paper or fee is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 on the date indicated above and is addressed to the Assistant Commissioner of Patents and Trademarks, Washington, D.C. 20231



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supported hydrogenation catalysts (WO 96/34896) enables what is almost complete hydrogenation to be achieved with a reduction of about 20 % in molecular weights. These catalysts comprise a special pore size distribution of the silica, which is characterised in that 98 % of the pore volume corresponds to a pore diameter larger than 600 Å. Said catalysts have specific surfaces between 14-17 m²/g and average pore diameters of 3800-3900 Å. Dilute solutions of polystyrene in cyclohexane, with a concentration ranging from 1% to a maximum of 8 %, are hydrogenated with degrees of hydrogenation greater than 98 % and less than 100 %.

- The examples described in the documents cited above indicate a reduction in the absolute molecular weight of hydrogenated polystyrene at polymer concentrations less than 2 %. In general, a reduction in molecular weight results in a deterioration in the mechanical properties of a hydrogenated polystyrene.
- The comparative example in WO 96/34896, in which a commercially available 5 % Rh/Al₂O₃ catalyst is used (Engelhard Corp., Beachwood, Ohio, USA) results in a degree of hydrogenation of 7 % and indicates the lower activity of an alumina support compared with a silica-supported catalyst.
- Surprisingly, it has now been found that aromatic polymers can be completely hydrogenated without a significant reduction in molecular weight, by using special catalysts, wherein the catalyst is defined in that 10 % at most of its pore volume corresponds to a pore diameter less than 600 Å, and it has an average pore diameter greater than 1000 Å and a specific surface greater than 3 m²/g, and comprises a defined pore size distribution.

The present invention relates to a process for the hydrogenation of aromatic polymers in the presence of catalysts, wherein a metal or a mixture of metals of subgroup VIII of the periodic table, together with a support comprising alumina, is used as a catalyst, and the pore volume corresponding to the pore diameter of the catalyst between 1000 and 10,000 Å, as measured by mercury porosimetry, is generally 100 - 15 %,

preferably 90 - 20 %, most preferably 80 - 25 %, particularly 70 to 30 %, with respect to the total pore volume as measured by mercury porosimetry.

The average pore diameter, as determined by mercury porosimetry, is greater than 1000 Å.

However, the mercury method is only satisfactory for pores which are larger than 60 Å. Pore diameters smaller than 600 Å are therefore determined by nitrogen sorption as described by Barret, Joyner and Halenda (DIN 66 134).

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The catalysts have a pore volume as measured by nitrogen sorption of less than 10 %, preferably less than 5 %, for pore diameters less than 600 Å. The pore volume as measured by nitrogen sorption is quoted with respect to the total pore volume as measured by mercury porosimetry.

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The average pore diameter and the pore size distribution are determined by mercury porosimetry according to DIN 66 133.

The average pore diameter generally ranges from 1000 Å to 10,000 Å, preferably from 2000 Å to 7000 Å, most preferably from 2500 Å to 6000 Å.

Methods for the characterisation of hydrogenation catalysts are described in WO 96/34896 (=US-A-5,612,422), and in Applied Heterogeneous Catalysis, Institute Français du Petrole Publication, pages 189-237 (1987), for example.

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The catalysts consist of metals of subgroup VIII, which are present together with a support comprising alumina.

Aluminas of general chemical formula Al₂O₃ occur in various modifications. A distinction is made, for example, between hexagonal α-Al₂O₃ and face-centred cubic crystalline γ-Al₂O₃. β-Al₂O₃ is understood to comprise a group of aluminas which contain small amounts of extraneous ions in their crystalline lattice. Other special

modifications exist, and numerous transitional forms exist between aluminium hydroxides and aluminas.

The specific surface of the catalyst is determined by the BET (Brunauer, Emmett and 5 Teller) method using nitrogen adsorption according to DIN 66 131 and DIN 66 132.

The (BET) specific surfaces for nitrogen are generally greater than $3 \text{ m}^2/\text{g}$, and preferably range from $5 \text{ m}^2/\text{g}$ to $80 \text{ m}^2/\text{g}$, most preferably from $8 \text{ m}^2/\text{g}$ to $60 \text{ m}^2/\text{g}$.

Metals of subgroup VIII are generally used, preferably nickel, platinum, ruthenium, rhodium, or palladium, most preferably platinum or palladium.

The metal content is generally 0.01 to 80 %, preferably 0.05 to 70 % with respect to the total weight of catalyst.

In a process which is operated as a batch process, the 50 % value of the cumulative distribution of particle size generally ranges from 0.1 μm to 200 μm , preferably from 1 μm to 100 μm , most preferably from 3 μm to 80 μm .

The usual solvents for hydrogenation reactions are aliphatic or cycloaliphatic hydrocarbons, aliphatic or cycloaliphatic saturated ethers, or mixtures thereof, e.g. cyclohexane, methylcyclopentane, methylcyclohexane, ethylcyclohexane, cyclooctane, cycloheptane, dodecane, dioxane, diethylene glycol dimethyl ether, tetrahydrofuran, isopentane and decahydronaphthalene.

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If aliphatic or cycloaliphatic hydrocarbons are used as solvents, they preferably contain water in a amount which ranges in general from 0.1 ppm to 500 ppm, preferably from 0.5 ppm to 200 ppm, most preferably from 1 ppm to 150 ppm, with respect to the total solvent.

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The process according to the invention generally results in what is practically the complete hydrogenation of the aromatic units. As a rule, the degree of hydrogenation

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is \geq 80 %, preferably \geq 90 %, most preferably \geq 99 % to 100 %. The degree of hydrogenation can be determined by NMR spectrometry or UV spectroscopy, for example. The process according to the invention most preferably results in hydrogenated aromatic polymers, particularly polyvinylcyclohexane, wherein the amount of dyads with a syndiotactic configuration is greater than 50.1 % and less than 74 %, particularly 52 to 70 %.

Aromatic polymers are used as the starting materials, and are selected, for example, from polystyrenes which are optionally substituted on their phenyl ring or on their vinyl group, or from copolymers thereof with monomers selected from the group comprising olefines, (meth)acrylates or mixtures thereof. Other suitable polymers include aromatic polyethers, particularly polyphenylene oxide, aromatic polycarbonates, aromatic polyesters, aromatic polyamides, polyphenylenes, polyphenylenes, polyphenylene vinylenes, polyphenylene ethinylenes, polyphenylene sulphides, polyaryl ether ketones, aromatic polysulphones, aromatic polyether sulphones, aromatic polyimides and mixtures thereof, and optionally copolymers with aliphatic compounds also.

Suitable substituents in the phenyl ring include C_1 - C_4 alkyl groups, such as methyl or ethyl, C_1 - C_4 alkoxy groups such as methoxy or ethoxy, and aromatic entities which are condensed thereon and which are bonded to the phenyl ring via a carbon atom or via two carbon atoms, comprising phenyl, biphenyl and naphthyl.

Suitable substituents on the vinyl group include C_1 - C_4 alkyl groups such as methyl, ethyl, or n- or iso-propyl, particularly methyl in the α -position.

Suitable olefinic comonomers include ethylene, propylene, isoprene, isobutylene, butadiene, cyclohexadiene, cyclohexene, cyclopentadiene, norbornenes which are optionally substituted, dicyclopentadienes which are optionally substituted, tetracyclododecenes which are optionally substituted, and dihydrocyclopentadienes which are substituted,

 C_1 - C_8 , preferably C_1 - C_4 alkyl esters of (meth)acrylic acid, preferably methyl and ethyl esters,

 C_1 - C_8 , preferably C_1 - C_4 alkyl ethers of vinyl alcohol, preferably methyl and ethyl 5 ethers,

C₁-C₈, preferably C₁-C₄ alkyl esters of vinyl alcohol, preferably vinyl acetate,

derivatives of maleic acid, preferably maleic anhydride, derivatives of acrylonitrile, preferably acrylonitrile and methacrylonitrile.

The preferred polymers are polystyrene, polymethylstyrene, and copolymers of styrene and at least one other monomer selected from the group consisting of α -methylstyrene, butadiene, isoprene, acrylonitrile, methyl acrylate, methyl methacrylate, maleic anhydride and olefines such as ethylene and propylene for example. Examples of suitable copolymers include those formed from acrylonitrile, butadiene and styrene, copolymers of acrylic esters, styrene and acrylonitrile, copolymers of styrene and α -methylstyrene, and copolymers of propylene, diene and styrene.

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The aromatic polymers generally have (weight average) molecular weights Mw from 1000 to 10,000,000, preferably from 60,000 to 1,000,000, most preferably from 70,000 to 600,000, particularly from 100,000 to 480,000, as determined by light scattering.

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The polymers can have a linear chain structure or can have branching locations due to co-units (e.g. graft copolymers). The branching centres may comprise star-shaped or branched polymers, or may comprise other geometric forms of the primary, secondary, tertiary or optionally of the quaternary polymer structure.

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The copolymers may exist as random copolymers or alternatively may exist as block copolymers.

Block copolymers comprise di-blocks, tri-blocks, multi-blocks and star-shaped block copolymers.

5 The initial polymers which are used are generally known (e.g. WO 94/21694).

The amount of catalyst to be used is described in WO 96/34896, for example.

The amount of catalyst used depends on the process which is carried out. This process can be conducted continuously, semi-continuously or batch-wise.

In a continuous system, the time of reaction is considerably shorter: it is influenced by the dimensions of the reaction vessel. In a continuous procedure, it is possible to use a trickling system or a liquid pool system, which both employ fixed catalysts, and it is also possible to use a system comprising a suspended catalyst, which can be recycled for example. Fixed catalysts can exist in the form of tablets or as an extruded product, for example.

The polymer concentrations with respect to the total weight of solvent and polymer generally range from 80 to 1, preferably from 50 to 10, particularly from 40 to 15 % by weight.

The reaction is generally conducted at temperatures between 0 and 500°C, preferably between 20 and 250°C, particularly between 60 and 200°C.

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The reaction is generally conducted at pressures from 1 bar to 1000 bar, preferably from 20 to 300 bar, particularly from 40 to 200 bar.

The palladium catalyst can be used either in reduced form or in unreduced form in the reaction concerned, whilst maintaining a comparable activity. In an industrial process, it is considerably more advantageous to employ the catalyst in unreduced form and not to have to carry out an additional, costly catalyst reduction step as in WO 96/34 896.

Examples:

The absolute (weight average) molecular weights Mw of the initial polymer and of the hydrogenated product were determined by light scattering. The absolute (number average) molecular weights Mn were determined by membrane osmosis. In Example 2, the relative Mw value obtained from the GPC determination (gel permeation chromatography using tetrahydrofuran as the elutant) compared with a polystyrene standard corresponded to the absolute molecular weights which were determined for the polystyrene used.

Examples 1 and 2

The catalysts used in the examples are characterised in Table 1.

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A 1 litre autoclave was flushed with inert gas. The polymer solution and optionally the unreduced catalyst were added (Table 2). After closing the autoclave, it was repeatedly pressurised with a protective gas and then with hydrogen. After releasing the pressure, the respective hydrogen pressure was set and the batch was heated with stirring to the corresponding reaction temperature. After the consumption of hydrogen had commenced, the reaction pressure was held constant.

The time of reaction was the time from heating up the batch until complete hydrogenation was effected of the polystyrene, or was the time until the consumption of hydrogen tended towards its saturation value.

After the reaction was complete, the polymer solution was filtered. The product was precipitated in methanol and dried. The isolated product had the physical properties listed in the Table.

Le A 33 510 - foreign countries

 Table 1:
 Physical characterisation of the catalysts used

	N (ata)	ivicial content					Č	%	0.5
	Total enerific	Surface (BETN ²	(177) 2000				# 5/2	м) М	11.9
	Average pore	diameter					₩		2722
	Pore volume for	pore diameters of	1000 A - 10,000 A	/ total mercury	pore volume	(38 A -15 µm)	%		63
	N ₂ pore volume	for pore diameters	of < 600 Å / total	mercury pore	volume		%		4
	Mercury pore	volume for pore	diameters from	1000 A - 10,000 A			mm³/g	700	400
	Total mercury	pore volume for	pore diameters	from	38 A - 15 µm		g/,wm	77.	ò
11.	Nitrogen pore	volume for pore	diameters of	< 600 Å		ŕ	mm/g	32	
. 14. July 19.	Catalyst INO.				0				

Catalyst: Engelhard De Meern, Holland, C 586-102, palladium on alumina

average pore diameter as measured by mercury porosimetry

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total specific nitrogen surface according to Brunauer, Emmett and Teller (BET, DIN 66131, DIN 66132)

Le A 33 510 - foreign countries

Fable 2.

Hydrogenation of polystyrene as a function of the catalyst, solvent system and reaction temperature

Example No.	Catalyst	Weight of	Solvent ³⁾	Weight of	Reaction	H ₂ pressure	Time of	Degree of	Tg	Mw
		polymer		catalyst	temperature		reaction	hydrogenation ⁴⁾	(DSC)	103
		ы	ם	as	ာ့	bar	min	%	ာ	lom/g
	5 % Pt/SiO ₂	2001)	2300	13.5	150	875 (psig)	110	98.4	146.6	
(comparative)		Mw ~	CH							
US-A		200,000	•							
5612422		g mol ^{-t}								
2	0.5 %	100.2 ²⁾	300 CYH	12.5	200	100	130	100	147	171
according to	Pd/Al ₂ O ₃ 5)					•			-	
the invention					:					

polystyrene (Mw ~ $200,000~g~mol^{-i}$)

polystyrene type 158 k; Mw = 280,000 g/mol, BASF AG, Ludwigshafen, Germany

CYH = cyclohexane

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determined by ¹H NMR spectrometry

Engelhard De Meern, Holland, palladium on alumina, C586-102

The platinum catalyst (Table 2), which was characterised in that 98 % of the pore volume corresponded to a pore diameter greater than 600 Å, did not completely hydrogenate polystyrene after 110 minutes at 150°C (98.4 %; comparative example 1).

5 The platinum catalyst according to Example 5 of US-A 5 612 422 resulted in a decrease of about 20 % in the molecular weight Mw.

A decrease in molecular weight results in particular in a reduction of the average molecular weight Mw (US-A 5 612 422). Surprisingly, hydrogenation according to the invention in the presence of the alumina catalyst resulted in no significant decrease in the absolute molecular weight Mw, even at elevated temperatures of reaction.

Within the range of accuracy of the measurements, the absolute average molecular weight Mw of the hydrogenated product corresponded to that of the polystyrene used.

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Compared with comparative example 1, the special alumina catalyst of the present invention resulted in a comparable time of reaction, but with complete hydrogenation being achieved at the same time, using a polystyrene of higher molecular weight and at a higher polymer concentration.

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The catalyst is distinguished by a weight of noble metal which is less by a factor of 10, which results in a significant reduction in raw material costs and which thus improves the economics of the hydrogenation process.

Le A 33 510 - foreign countries

- 12 -

<u>Claims</u>

1. A process for the hydrogenation of aromatic polymers in the presence of catalysts, wherein a metal or a mixture of metals of subgroup VIII of the periodic table, together with a support comprising alumina, is used as a catalyst, and the pore volume corresponding to the pore diameter of the catalyst between 1000 and 10,000 Å, as measured by mercury porosimetry, is generally 100 - 15 % with respect to the total pore volume as measured by mercury porosimetry.

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- 2. A process according to claim 1, wherein the pore volume is 90 to 20 % with respect to the total pore volume.
- 3. A process according to claim 2, wherein the pore volume is 80 to 25 % with respect to the total pore volume.
 - 4. A process according to claim 3, wherein the pore volume is 70 to 30 % with respect to the total pore volume.
- 20 5. A process according to one or more of the preceding claims, wherein the catalysts have a pore volume, as measured by nitrogen sorption, of less than 10 % for pore diameters less than 600 Å.
- 6. A process according to one or more of the preceding claims, wherein the metals are selected from nickel, platinum, ruthenium, rhodium and palladium.
 - 7. A process according to one or more of the preceding claims, wherein the process is conducted in the presence of solvents which can be used for hydrogenation reactions.

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8. A process according to one or more the preceding claims, wherein the catalyst has a specific surface greater than 5 m²/g.

Le A 33 510 – foreign countries

- 13 -

A process for the hydrogenation of aromatic polymers in the presence of special catalysts

Abstract

A process for the hydrogenation of aromatic polymers in the presence of catalysts, wherein a metal or a mixture of metals of subgroup VIII of the periodic table, together with a support comprising alumina, is used as a catalyst, and the pore volume corresponding to the pore diameter of the catalyst between 1000 and 10,000 Å, as measured by mercury porosimetry, is generally 100 - 15 % with respect to the total pore volume as measured by mercury porosimetry.

COMBINED DECLARATION AND POWER OF ATTORNEY

ATTORNEY DOCKET NO

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name. I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought

on the invention entitled

METHOD FOR HYDROGENATING AROMATIC POLYMERS IN THE PRESENCE OF SPECIAL CATALYSTS

the specification of which is attached hereto,

or was filed on January 26, 2000

as a PCT Application Serial No. PCT/EP00/00576

I hereby state that I have reviewed and understand the contents of the aboveidentified specification, including the claims.

I acknowledge the duty to disclose information which is material to the patentability of this application in accordance with Title 37, Code of Federal Regulations, §1.56.

I hereby claim foreign priority benefits under Title 35, United States Code, \$119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:

Prior Foreign Application(s), the priority(ies) of which is/are to be claimed:

199 04 612.3 (Number)

Germany (Country)

February 5, 1999 (Month/Day/Year Filed)

AUG 3 0 2001

I hereby claim the benefit under Title 35, United States Code, \$120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, \$112, I acknowledge the duty to disclose the material information as defined in Title 37, Code of Federal Regulations, \$1.56 which occured between the filing date of the prior application and the national or PCT international filing date of this application:

(Application Serial No.) (Filing Date) (Status)

(Application Serial No.) (Filing Date) (Status)

(Application Serial No.) (Filing Date) (Status)

(patented, pending, abandoned)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment,

ful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Le A 33 510-US

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